Signaling pathways involved in apoptosis induced by novel angucycline antibiotic landomycin E in Jurkat T-leukemia cells

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Aim. To study the molecular mechanisms of action of novel anticancer antibiotic landomycin E (LE). Methods. Annexin V/propidium iodide, DAPI (4',6-diamidino-2-phenylindole) staining, Western-blot analysis. Results. LE applied in 2 µg/ml dose (IC50), induced reactive oxygen species (ROS)-dependent splitting of poly [ADP-ribose] polymerase 1 (PARP-1) and DNA Fragmentation Factor 45 (DFF45) proteins involved in DNA reparation. This effect was observed 6 h after the start of treatment and it positively correlated with phosphatidyl serine externalization (early morphological marker of apoptosis). We suggest that cleavage of PARP-1 and DFF45 was mediated by active caspase-7 which is a key effector caspase in the LE-induced apoptosis in leukemia cells. We found that activation of initiator procaspase-10 (involved in receptor-mediated apoptosis) was the earliest detected event in LE-induced apoptotic signaling pathways; however, this activation was shown to be ROS-independent. We also demonstrated that the induction of apoptosis by LE is accompanied by activation of apoptosis-inducing factor (AIF) in mitochondria. Conclusions. Our data suggest that LE-induced cascade of apoptotic events is started by the initiator caspase-10 which leads to activation of the effector caspase-7 and AIF that is known to induce caspase-independent apoptosis involving ROS generation.

Keywords: tumor cells, landomycin E, apoptosis, caspases, AIF, reactive oxygen species.

Introduction. Streptomyces species play a significant role in the production of bioactive natural products, many of which are polyketides, such as anthracyclines and tetracyclines [1]. The angucyclines are the largest group of polycyclic aromatic polyketides with more than 120 members that is constantly growing [2, 3]. The most characteristic feature of angucyclines is their uniquely shaped benz[a]anthracene tetracyclic framework with an angularly condensed ring [4]. The group is rich in chemical scaffolds and various biological activities, predominantly antitumor and antibacterial. Yet, none of these compounds have been developed to clinically applied drugs usually due to toxicity or solubility issues [1–3]. Landomycins are the most perspective group of angucyclines possessing strong anti-neoplastic potential.

All natural landomycins identified to date share the same aglycon (landomycinone) and vary only in their oligosaccharide chain, a linear glycosidic chain containing only di- and trideoxysugars (β-D-olivose and α-L-rhodinose) [5]. They show broad activity against many cancer cell lines with the general tendency that
compounds with longer saccharide chains show higher activity [6, 7].

The main compound, landomycin A, containing a hexasaccharide side chain, has so far been shown to be the most potent congener and was extensively tested by the National Cancer Institute (USA) towards 60 selected human cancer cell line panel and particularly towards prostate cancer lines [8, 9].

Landomycin E (LE) is a novel representative of landomycins synthesized by *Streptomyces globisporus* strain 1912 growing in a soybean culture medium [10]. It contains 3 saccharide residues (α-L-rhodinose-(1 → 3)-β-D-olivose-(1 → 4)-β-D-olivose) conjugated to an angular tetracyclic quinone. Antitumor action of LE was demonstrated against various tumor cell lines in vitro [11] and Guerin carcinoma in rats in vivo [12]. It was also reported that LE is able to overcome resistance to doxorubicin, vincristin and colchicine in cancer cells, overexpressing various types of ABC-transporters (P-gp, MRP-1, bcrp) [13, 14]. However, molecular mechanisms of its antineoplastic effects still require clarification.

Flow cytometry experiments showed that landomycin A specifically blocked cell cycle progression from G1 phase to S phase (DNA synthesis) [8]. Structurally unrelated anticancer drugs (e. g. bleomycin, mitomycin C, and neocarzinostatin) possess a similar pattern of cell cycle inhibition. However, in contrast to many clinically useful drugs of a similar structure, like the anthracyclines and chromomycins, landomycins do not bind directly to DNA [4, 13, 14]. Landomycin E whose glycosidic chain contains only 3 deoxysugars, does not possess cell cycle specificity [14], but it strongly impairs mitochondria functions leading to the generation of reactive oxygen species (ROS). Nevertheless, role of mitochondria and ROS in LE-mediated apoptosis remains poorly understood. It needs to be explained whether the LE-induced early generation of ROS in mitochondria can trigger caspase activation, or it is only a supplementary step in general scheme of apoptosis induced by this drug.

The main goal of present study was to investigate in more detail specific apoptotic signaling pathways which are induced by the landomycin E in leukemia cells. This might allow to identify potential molecular targets of LE action in tumor cells.

Materials and methods. Landomycin E (95 % purity, according to TLC data) was prepared in the laboratory of Dr. B. Matselyukh at D. K. Zabolotny Institute of Microbiology and Virology, National Academy of Sciences of Ukraine. Doxorubicin produced by «Ebeve» (Austria) was bought at the local pharmacy.

Human Jurkat T-leukemia cells were obtained from cell culture collections at R. E. Kavetsky Institute of Experimental Pathology, Oncology and Radiobiology, National Academy of Sciences of Ukraine. Cells were cultured in RPMI 1640 medium supplemented with 10 % fetal calf serum («Sigma Chemical Co.», USA), 50 µg/ml streptomycin («Sigma Chemical Co.»), 50 units/ml penicillin («Sigma Chemical Co.») in 5 % CO₂-containing humidified atmosphere at 37°C.

For experiments, cells were seeded into 24-well tissue culture plates («Greiner Bio-one», Germany). The cytotoxic effect of antitumor drugs was studied under the inverted microscope («Biolam-P1», «LOMO», Russian Federation) after cell staining with trypan blue dye (0.1 %). N-acetylcysteine («Sigma», USA) was dissolved in 1 ´ phosphate buffered saline (PBS) and added to cell culture 30 min before addition of anticancer drugs (final concentration 1 mM). Broad caspase inhibitor z-VAD-fmk («BD Pharmingen», USA), 20 mM stock solution in DMSO was dissolved in cell culture medium prior to addition to cell culture to achieve final concentration 100 µM.

FITC-conjugated annexin V («BD Pharmingen», USA) and propidium iodide («Sigma») double staining was performed to detect early apoptotic events under treatment of Jurkat cells by LE. At 1, 3, 6, 12, 24 h after addition of LE Jurkat cells were centrifuged at 2000 rpm, washed twice with 1 ´ PBS, and incubated for 15 min in annexin V binding buffer («BD Pharmingen»), 20 mM stock solution in DMSO was dissolved in cell culture medium prior to addition to cell culture to achieve final concentration 100 µM.

FITC-conjugated annexin V («BD Pharmingen», USA) and propidium iodide («Sigma») double staining was performed to detect early apoptotic events under treatment of Jurkat cells by LE. At 1, 3, 6, 12, 24 h after addition of LE Jurkat cells were centrifuged at 2000 rpm, washed twice with 1 ´ PBS, and incubated for 15 min in annexin V binding buffer («BD Pharmingen») containing 1/20 volume of FITC-conjugated annexin V solution and PI (20 µg/ml). 10 µl of cell suspension was placed on slides.

Cytomorphological investigations were performed using Zeiss Axiosmager A1 fluorescent microscope («Zeiss», Germany).

DAPI staining was performed to study chromatin condensation in Jurkat cells at various timepoints of LE treatment. 1, 3, 6, 12, and 24 h after the addition of LE, Jurkat cells were centrifuged at 2,000 rpm, washed twice with 1 ´ PBS, fixed in 4 % paraformaldehyde so-
solution for 15 min at room temperature, and then permeabilized by 0.1% Triton X-100 solution in PBS for 3 min. After that, cells were incubated with 1 µg/ml DAPI solution (4',6-diamidino-2-phenylindole) («Sigma») for 5 min, washed twice with PBS, 10 µl of cell suspension was added to slides and cover glasses placed. Cytomorphological investigations were performed on Zeiss AxioImager A1 fluorescent microscope.

Western blot analysis was used to evaluate expression of proteins, involved in signaling pathways of apoptosis, induced by landomycin E, and performed, as described before [15]. Membrane was incubated with monoclonal rabbit antibodies raised against cleaved caspase-3, cleaved caspase-6, cleaved caspase-7, Bid («Cell Signaling», USA), AIF (sc-5586), Bax (sc-6236), Bcl-2 (sc-492), Bcl-X_{L/S} (sc-634) («Santa Cruz Biotech», USA), caspase-9 (BD 556585) («BD Pharmingen»), β-actin («Sigma»), monoclonal goat antibodies against procaspase-3 (sc-1226) («Santa Cruz Biotech») and monoclonal mouse antibodies against caspase-2, caspase-8 («BD Pharmingen»), caspase-10 («MBL», USA) for 12 h at 4°C at slow shaking. Dilution of primary antibodies was 1:1,000 in 5% BSA, 0.1% PBS-Tween, except the antibodies against β-actin diluted (1:5000) as recommended by supplier.

**Results and discussion.** LE was shown to be highly active for all studied tumor cell lines of both epithelial and mesenchymal origin, and its effect was comparable with that of doxorubicin and vincristin. Leukemia cells were most sensitive to LE action [11]. That is why we used human T-leukemia Jurkat cells in our further studies.

Phosphatidylserine translocation to the external layer of the cell membrane of dying cells measured by annexin V test is considered to be one of the earliest hallmarks of apoptosis [16]. Landomycin E (2 µg/ml) induced early phosphatidylserine translocation already at 6 h after treatment of Jurkat cells, while doxorubicin in the same concentration caused PS exposure only at 12 h after the treatment start (Fig. 1, A, see inset). Massive DNA fragmentation was measured by cytomorphological evaluation of apoptotic bodies and hypercondensed chromatine in Jurkat cells using DAPI staining. Such fragmentation has appeared only at 12 h both after LE and Dx treatment (Fig. 1, B, see inset). Thus, LE induces more rapid apoptosis, than Dx, in spite of LE’s lower cytotoxic activity [11, 13, 14]. Doxorubicin is classified as a topoisomerase II poison [17], although other mechanisms of its action have been also proposed: ROS generation, DNA intercalation, inhibition of nucleic acid synthesis [18, 19]. Thus, an induction of apoptosis by Dx can be explained by its DNA-damaging nature, while LE may use other pathways taking place more rapidly than at DNA damaging in p53-dependent apoptosis.

Mitochondria are perfect candidate for LE targeting. ROS are overproduced by mitochondria under pathological conditions, such as ionizing radiation, heat shock or anticancer drug action [20]. ROS scavenger N-acetylcysteine (NAC) was used to study in detail ROS involvement in LE and Dx apoptotic signaling in Jurkat cells.

Pre-treatment of Jurkat cells for 30 min with 1 mM NAC restored number of live cells to control levels at 6 h and 12 h after LE treatment (P < 0.01), while at 24 h no significant effect of NAC at IC50 concentration of LE (2 µg/ml) could be observed (Fig. 2). It is known, that Dx anticancer activity also involves ROS [18], but in our case, no statistically significant differences between Dx-treated and Dx + NAC-pre-treated cells were found (Fig. 2).

Caspase-dependent signaling pathways were studied by Western-blot analysis.

Activation of key effector caspases – caspase-3 which cleaves enzymes PARP-1 and DFF45, as well as activation of caspase-6 which cleaves nuclear lamina, take place only at 24 h after LE treatment of Jurkat cells (Fig. 3). This was also confirmed by a study of expression of procaspase-3 that was cleaved only at 24 h, while at all time points from 1h till 12 h no signs of its cleavage were detected. However, slight activation of other key effector caspase-7 was observed already at 6 h after LE treatment of Jurkat cells, which perfectly correlated with time-dependent cleavage of PARP-1 and DFF45 (Fig. 3).

In contrast to LE, doxorubicin (2 µg/ml) induced simultaneous activation of effector caspases-3 and -7 at 12 h, while active form of effector caspase-6 appeared only at 24 h after treatment (Fig. 4). We suggest that caspase-7 plays a key role in late stages of LE-induced apoptosis in leukemia cells, while caspase-3 and caspase-6 are not so important.
Pre-treatment of Jurkat cells with 1 mM NAC for 30 min led to inhibition of production of reactive oxygen species under LE treatment and also completely blocked activation of caspase-7 and cleavage of DFF45 and PARP-1 by this enzyme at 6, 12 h, and partly at 24 h (Fig. 3).

However, there were no significant changes in the levels of proapoptotic protein Bax involved in mitochondria-induced apoptosis (Fig. 5). Thus, one can see that ROS do not activate Bax and, thus, their role in LE-induced apoptosis is supplementary, not initiatory. This can also be confirmed by absence of changes in expression of antiapoptotic proteins of Bcl-2 family – Bcl-XL and Bcl-2 itself (Fig. 5), which clearly indicates that small mitochondrial proteins are not involved in LE-induced apoptosis. NAC pre-treatment also had no effect on their levels in target cells. Also in case of LE cleavage of caspase-9 (key initiator caspase, involved in mitochondria-mediated apoptosis) takes place only at 24 h after treatment (Fig. 5) and cytochrome C release from mitochondria – only at 12 h (data not shown), while effector caspase-7 was activated already at 6 h after LE treatment. This again indicates important, but not initiatory role of mitochondria in these processes.

That is why FAS/CD95 receptor in plasma membrane of cells can be another potential target for this drug [21]. Since all landomycins have linear glycosidic chain, they can interact in some way with extracellular part of CD95 receptor or cell glycocalyx and induce receptor-mediated apoptosis via caspase-8 or caspase-10. This suggestion can be indirectly supported by the results of National Cancer Institute testing of landomycin A, landomycin E and landomycin D, where landomycins with longer glycosidic chain had significantly higher antineoplastic activity (see www.dtp.nci.nih.gov). To study this hypothesis, Western-blot analysis was performed with antibodies against initiator caspases-2, -8, -10.

It was found that LE (2 µg/ml, 24 h) induced cleavage of all tested initiator caspases, but at early stages (6 h), caspase-10 was the only active enzyme, and its activation was not blocked by NAC (Fig. 5). Thus, caspase-10 is the only initiator enzyme, which is activated at earliest timepoint by LE, and its activation is not ROS-mediated. So, caspase-10 seems to act upstream of mitochondria in the LE-induced apoptosis.

Doxorubicin is also known to induce ROS production in target cells [18], but this effect was much weaker than that of LE (see Fig. 2). NAC pre-treatment...
had no effect in diminishing cytotoxicity of Dx \textit{in vitro} in Jurkat cells, but partly inhibited activation of effector caspases-3, -7 and their cleavage of PARP-1 and DFF45 at 12 h after Dx treatment of Jurkat cells (see Fig. 4). No effect of NAC was observed at 24 h time-point, which indicates subsidiary role of ROS in Dx-induced apoptosis. It is known that Dx activates mitochondrial pathway of apoptosis via p53 protein or, when it is absent (as in case with Jurkat cells where p53 is inactivated) via cleavage of small mitochondrial Bid protein via caspase-8 [22], which was also observed in our case (Fig. 5). Landomycin E led to Bid cleavage only in 24 h which excludes caspase-8-Bid bridge in apoptosis induction. This suggests principal difference of the mechanisms of LE-induced cell death from Dx-induced ones.

Most anticancer drugs induce classical caspase-dependent apoptosis. However, several chemotherapeutic drugs of clinical use and some natural agents with anti-tumorigenic properties can also lead to caspase-independent type of cell death potentiated by specific «backup death pathways» [23]. Broad caspase inhibitor z-VAD-fmk was used to find out what death pathway is preferentially stimulated by LE in Jurkat cells. Pre-treatment of Jurkat cells for 1 h with 100 μM of z-VAD-fmk attenuated Dx-induced (2 μg/ml) apoptosis for 24 h, but failed to stop LE-induced (2 μg/ml) apoptosis measured by counting annexin V + /PI-cells (Fig. 6).

Apoptosis-inducing factor (AIF) whose level was significantly increased during LE-induced apoptosis (Fig. 5), can be another candidate for key apoptotic mediator.

AIF was the first mitochondrial protein shown to mediate caspase-independent cell death [24]. It was initially characterized as a protein confined within the mitochondrial intermembrane space of healthy cells. During apoptosis AIF is released from mitochondria and translocates to the nucleus, where it mediates nuclear features of apoptosis such as chromatin condensation and large-scale (~50 kb) DNA degradation [25]. In healthy cells, AIF plays role in oxidative phosphorylation and redox control [26]. Thus, AIF can generate ROS playing role downstream in the LE-induced signaling apoptotic pathways. We found a significant increase in AIF level in Jurkat cells treated with LE (Fig. 4), which confirms our suggestion of its role in LE action.

Thus, it is AIF, but not caspase-7, which seems to play a major role at terminal stages of LE-induced apoptosis. This might explain why NAC, even used for 24 h, does not stop death of LE-treated cells. NAC can block activation of effector caspases, but not of AIF, which itself is a source of ROS, and thus NAC pre-treatment cannot stop AIF’s proapoptotic action.

In conclusion, despite similar chemical structure, doxorubicin and landomycin E use different signaling pathways inducing apoptosis. Dx-induced apoptosis takes place later than LE-induced one, and its first signs are observed only at 12 h after Dx treatment of Jurkat cells. Dx induces apoptosis via DNA damage and mito-
chondrial pathway, confirmed by cleavage of initiator procaspase-8, its substrate Bid and procaspase-9 leading to subsequent downstream activation of effector members of caspase cascade. In contrast to Dx, LE may use CD95 receptor to activate cleavage of caspase-10 that occurs 6 h after LE treatment and is ROS-in-
dependent process which is not blocked by NAC. Then caspase-10 can cleave caspase-7 (6 h) which activates mitochondria-signaling pathway accompanied by a release of AIF (6 h), ROS generation, and cleavage of PARP-1 and DFF45 (6 h). Since broad caspase inhibitor z-VAD-fmk does not block apoptosis induced by LE even after 24 h treatment, we suggest that effector caspases play a supplementary role at late stages of LE-induced apoptosis, while AIF is a key effector protein that orchestrates apoptosis network upon LE action.

Conclusions. LE, despite lower cytotoxic activity comparing with Dx, induces more rapid development of apoptosis. This suggests different molecular mechanisms of the cytotoxic action of LE. The results of Western-blot analysis indicate that LE induces apoptosis of mixed type. At early stages of LE action, the initiator caspase-10 involved in the receptor-mediated apoptosis is important, while at later stages, AIF release from mitochondria takes place, and this leads to ROS generation and caspase-independent cleavage of DNA that is confirmed by caspase inhibition tests. Such dual LE action suggests that this antibiotic could be used as an effective anticancer drug of second line of defense. This predicts LE ability to overcome tumor refractoriness to treatment with anticancer drugs (e.g., doxorubicin, cisplatin, vincristin) killing cancer cells via classic caspase-dependent apoptotic mechanisms.

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P. R. Panchuk
Сигнальні шляхи, залучені до індукації апоптозу новітнім антицикліновим антимикробіом Ланодіміцином Е в клітинах T-лейкемії людини лінії Jurkat

Резюме
Мета. Дослідити молекулярні механізми дії нового протицинкілінового антибіотика Ланодіміцину Е (ЛЕ). Методи. Фарбування клітин анексином V, йодистим пропідіїсм, DAPI (4',6-діаміно-2-феніліндоз), Вестерн-блот аналіз. Результати. Визначено, що ЛЕ (2 мкг/мл) індукує розщеплення білків PARP-1 (полі [ADP-рибоза] полімераза 1) та DFF45 (DNA Fragmentation Factor 45), залучених до репарації ДНК, вже через 6 год після додавання до клітинної культури. Це явно опосередковане активними сполуками кисню (АСК) і позитивно корелює з екстремізацією фосфатидилхоліну на поверхні плазматичної мембрани (ранній маркер апоптозу). Розщеплення PARP-1 і DFF45 здійснюється активною каспазою-7 – ключовою ефекторною каспазою у ЛЕ-опосередкованому апоптозі в лейкемічних клітинах. Найбільшою подією у сигнальних шляхах апоптозу, спричиненого ЛЕ, є виявлення активації ініціаторної каспази-10, причиною до рецептор-опосередкованого апоптозу. Показано, що цей процес є АСК-незалежним. Нами також встановлено, що індукація апоптозу ЛЕ супроводжується ак- тівацією апоптоз-індукуючого фактора (АПФ) в мітохонд- ріях. Висновки. ЛЕ-індукуваний каскад апоптичних подій по- чинається з активації ініціаторної каспази-10, що приводить до подальшої активації ефекторної каспази-7 та АФ, який здатний індукувати каспаз-незалежний апоптоз за участі АСК.

Ключові слова: зовнішні клітини, ланодіміцин, апоптоз, каспази, апоптоз-індукуючий фактор, активні сполуки кисню.

P. R. Panchuk
Сигнальні пути, участиючі в індукації апоптоза новим антицикліновим антимикробіом Ланодіміцином Е в клітинах T-лейкемії людини лінії Jurkat

Резюме
Цель. Исследовать молекулярные механизмы действия нового протицинкілінового антибіотика ланодіміцину Е (ЛЕ). Методы. Окрашивание клеток анексином V, йодистым пропідіїсм, DAPI (4',6-діаміно-2-феніліндоз), Вестерн-блот анализ. Результаты. Определено, что ЛЕ (2 мкг/мл) индукует расщепление белков PARP-1 (полі [ADP-рибоза] полімераза 1) та DFF45 (DNA Fragmentation Factor 45), причастных к репарации ДНК, уже через 6 ч после добавления к клеточной культуре. Это явление опосредовано активными соединениями кислорода (АСК) и положительно коррелирует с экстремизацией фосфатидилхоллина на поверхности плазматической мембраны (ранний маркер апоптоза). Расщепление PARP-1 и DFF45 осуществляется активной каспазой-7 – ключевой эффекторной каспазой в
APOTOPSIS INDUCED BY ANTIBIOTIC LANDOMYCIN E IN JURKAT T LEUKEMIA CELLS

LE-oposerdemomom в лейкемичых клетках. Наиболее ранним событием в сигнальных путях апоптоза, вызванного ЛЕ, оказалась активация инциаторной прокаспазы-10, участвующей в рецептор-oposердемомом апоптозе. Показано, что этот процесс является АСК-независимым. Нами также установлено, что индукция апоптоза ЛЕ сопровождается активацией апоптоз-индуцирующего фактора (АИФ) в митохондриях.

Выводы. ЛЕ-индуцированный каскад апоптотических событий начинается с активации инциаторной каспазы-10, что приводит к дальнейшей активации эффекторной каспазы-7 и АИФ, который способен индуцировать каспазо-независимый апоптоз при участии апоптотических форм клетки.

Ключевые слова: эозиноположительные клетки, лейкемия Е, апоптоз, каспазы, апоптоз-индуктирующий фактор, активные соединения клетки.

REFERENCE


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